Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG112321\

Data File : BG051196.D

Acq On : 24 Nov 2021 00:02

Operator : CG/JU Sample : M4780-03MSD

Misc

ALS Vial : 17 Sample Multiplier: 1

Quant Time: Nov 24 06:56:42 2021

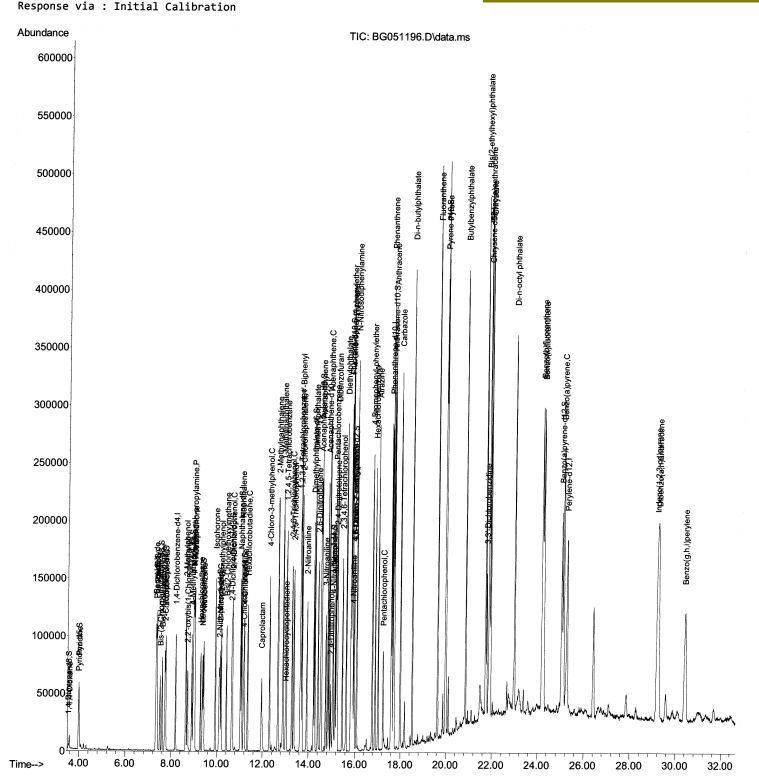
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\SFAM-EPA-BG112321.M

Quant Title : SVOA CALIBRATION
QLast Update : Wed Nov 24 06:04:50 2021

Instrument : BNA\_G ClientSampleId : DBLQ4MSD

## **Manual IntegrationsAPPROVED**

Reviewed By :Jagrut Upadhyay 11/24/2021 Supervised By :mohammad ahmed 11/30/2021



#### Quantitation Report (Qedit)

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG112321\

Data File : BG051196.D

Acq On : 24 Nov 2021 00:02

Operator : CG/JU Sample : M4780-03MSD

Misc

ALS Vial : 17 Sample Multiplier: 1

Quant Time: Nov 24 06:56:42 2021

 $\label{lem:quant_method} {\tt Quant_Methods\SFAM-EPA-BG112321.M}$ 

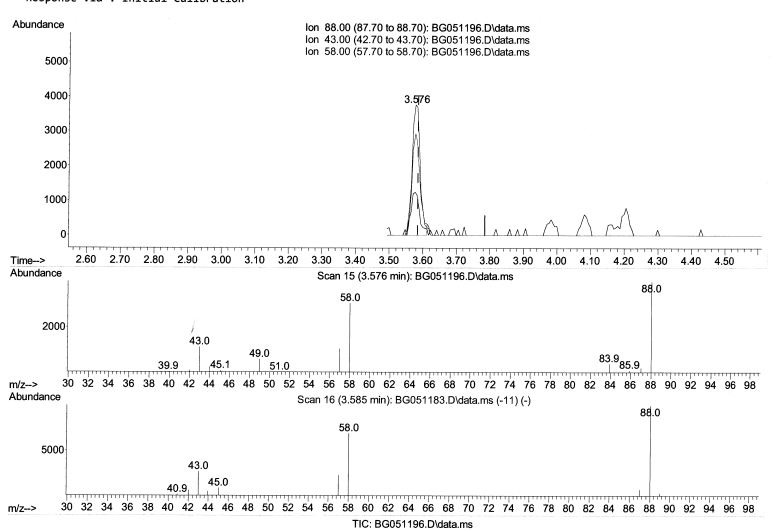
Quant Title : SVOA CALIBRATION

QLast Update : Wed Nov 24 06:04:50 2021 Response via : Initial Calibration



## **Manual IntegrationsAPPROVED**

Reviewed By :Jagrut Upadhyay 11/24/2021 Supervised By :mohammad ahmed 11/30/2021



### (2) 1,4-Dioxane

3.576min (-0.009) 6.84 ng/uL

response	6329	
Ion	Ехр%	Act%
88.00	100.00	100.00
43.00	28.70	32.94
58.00	78.00	77.87
0.00	0.00	0.00

### Quantitation Report (Qedit)

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG112321\

Data File : BG051196.D

Acq On : 24 Nov 2021 00:02

Operator : CG/JU

Sample : M4780-03MSD

Misc

ALS Vial : 17 Sample Multiplier: 1

Quant Time: Nov 24 06:56:42 2021

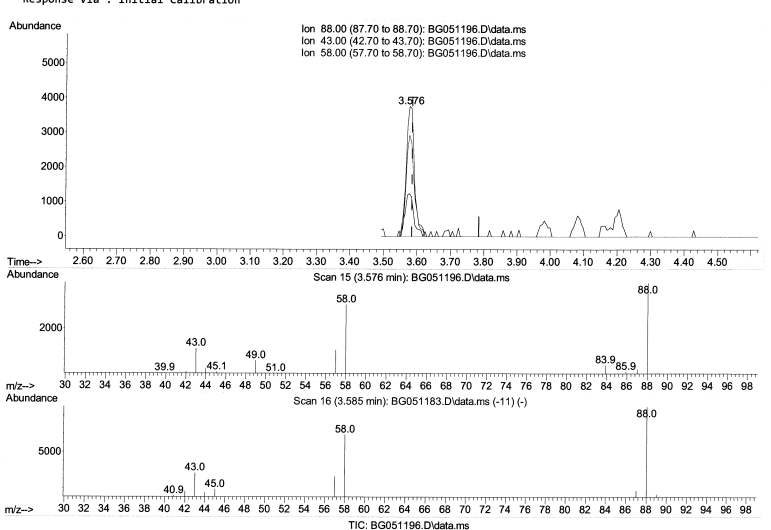
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\SFAM-EPA-BG112321.M

Quant Title : SVOA CALIBRATION

QLast Update : Wed Nov 24 06:04:50 2021 Response via : Initial Calibration Instrument : BNA\_G ClientSampleld : DBLQ4MSD

## Manual IntegrationsAPPROVED

Reviewed By :Jagrut Upadhyay 11/24/2021 Supervised By :mohammad ahmed 11/30/2021



#### (2) 1,4-Dioxane

response	6420		
Ion	Ехр%	Act%	
88.00	100.00	100.00	
43.00	28.70	32.94	
58.00	78.00	77.87	
0.00	0.00	0.00	

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG112321\

Data File : BG051196.D

Acq On : 24 Nov 2021 00:02 Operator : CG/JU Sample : M4780-03MSD

Misc

ALS Vial : 17 Sample Multiplier: 1

Quant Time: Nov 24 06:56:42 2021

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\SFAM-EPA-BG112321.M

Quant Title : SVOA CALIBRATION

QLast Update : Wed Nov 24 06:04:50 2021 Response via : Initial Calibration

**Instrument :** BNA\_G ClientSampleId : DBLQ4MSD

# **Manual IntegrationsAPPROVED**

Reviewed By :Jagrut Upadhyay 11/24/2021 Supervised By:mohammad ahmed 11/30/2021

	Compound	R.T.	QIon	Response	Conc Un:	its Dev	(Min)	
Inte	rnal Standards							
1)	1,4-Dichlorobenzene-d4	8.200	152	28507	20.000	ng/ul	0.00	
	Naphthalene-d8	11.026		127956	20.000		0.00	
	Acenaphthene-d10	14.828		84166	20.000	-	0.00	
	Phenanthrene-d10	17.577		178805	20.000		0.00	
	Chrysene-d12	21.878		156410	20.000		0.00	
	Perylene-d12	25.274		155720	20.000		0.00	
Syst	em Monitoring Compounds							
3)	1,4-Dioxane-d8	3.541	96	2091	2.549	ng/uL	0.00	
4)	Pyridine-d5	3.976	84	19271	8.006		0.00	
7)	Phenol-d5	7.354	99	51677	18.342		0.00	
9)	Bis-(2-Chloroethyl)eth	7.513	67	32497	18.365		0.00	
	2-Chlorophenol-d4	7.730		38586	19.019	-	0.00	
	4-Methylphenol-d8	8.911	113	39720	17.470	_	0.00	
	Nitrobenzene-d5	9.375		20338	18.829	-	0.00	
	2-Nitrophenol-d4	10.104		22666	18.603		0.00	
	2,4-Dichlorophenol-d3	10.644		41727	20.184		0.00	
	4-Chloroaniline-d4	11.161	131	42427	14.026		0.00	
	Dimethylphthalate-d6	14.222		133943	20.683	-	0.00	
	Acenaphthylene-d8	14.528		167928	20.564		0.00	
	4-Nitrophenol-d4	15.045		19310	18.421	_	0.00	
	Fluorene-d10	15.821	176	123180	21.122	-	0.00	
	4,6-Dinitro-2-methylph			23651	21.436		0.00	
	Anthracene-d10	17.677		187893	21.972	_	0.00	
•	Pyrene-d10	19.957		213928	22.604		0.00	
	Benzo(a)pyrene-d12	25.045	264	184519	22.187		0.00	
Targe	et Compounds					Ova	alue	
2)	1,4-Dioxane	3.576	88	6420m	> 6.939	-		74
	Pyridine	3.993	79	32118	12.823	_	99	•
	Benzaldehyde	7.336	77	37754	21.042		94	
	Phenol	7.383	94	67096	22.988		98	
10)	Bis(2-Chloroethyl)ether	7.607	93	46007	20.835	-	97	
	2-Chlorophenol	7.759	128	45125	21.826		96	
	2-Methylphenol	8.647	108	44430	20.436	-	98	
14)	2,2'-oxybis(1-Chloropr	8.711	45	67626	21.223		99	
	Acetophenone	9.028	105	74188	21.096		99	
	N-Nitroso-di-n-propyla	8.999	70	42589	21.074	-	99	
	4-Methylphenol	8.976	108	50373	21.668	-	93	
	Hexachloroethane	9.287	117	18317	20.975		95	
22)	Nitrobenzene	9.416	77	60784	21.461		95	
	Isophorone	9.933	82	119083	21.641		99	
	2-Nitrophenol	10.133	139	27405	21.715		99	
	2,4-Dimethylphenol	10.180	107	43644	16.914	•	96	
	Bis(2-Chloroethoxy)met	10.415	93	65407	21.532	•	99	
	2,4-Dichlorophenol	10.674	162	45821	22.516	_	99	
	Naphthalene	11.079	128	151129	21.707		98	
	4-Chloroaniline	11.191	127	53108	17.489	_	98	
	Hexachlorobutadiene	11.343	225	30112	21.453		96	
•	Caprolactam	11.949	113	17276	21.594	_	95	
	4-Chloro-3-methylphenol	12.301	107	58040	23.743	_	98	
•	<b>,</b> , –					J		

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG112321\

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Misc

ALS Vial : 17 Sample Multiplier: 1

Quant Time: Nov 24 06:56:42 2021

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\SFAM-EPA-BG112321.M

Quant Title : SVOA CALIBRATION QLast Update : Wed Nov 24 06:04:50 2021 Response via : Initial Calibration **Instrument :** BNA\_G

ClientSampleId: DBLQ4MSD

## **Manual IntegrationsAPPROVED**

Reviewed By: Jagrut Upadhyay 11/24/2021 Supervised By: mohammad ahmed 11/30/2021

Compound	R.T.	QIon	Response	Conc Units Dev(	(Min)
36) 2-Methylnaphthalene	12.671	142	105787	22.338 ng/ul	98
37) 1-Methylnaphthalene	12.889	142	105161	21.584 ng/ul	97
39) 1,2,4,5-Tetrachloroben	13.030	216	59985	22.702 ng/ul	96
40) Hexachlorocyclopentadiene	13.000	237	8580	8.034 ng/ul	94
41) 2,4,6-Trichlorophenol	13.271	196	40450	24.395 ng/ul	96
42) 2,4,5-Trichlorophenol	13.353	196	45922	26.447 ng/ul	95
43) 1,1'-Biphenyl	13.664	154	141667	22.536 ng/ul	98
44) 2-Chloronaphthalene	13.717	162	113637	22.725 ng/ul	99
45) 2-Nitroaniline	13.917	65	41903	24.212 ng/ul	94
47) Dimethylphthalate	14.269	163	151514	23.114 ng/ul	99
48) 2,6-Dinitrotoluene	14.405	165	33182	24.099 ng/ul	93
50) Acenaphthylene	14.557	152	185309	22.968 ng/ul	97
51) 3-Nitroaniline	14.739	138	30659	22.526 ng/ul	99
52) Acenaphthene	14.892	153	123592	23.228 ng/ul	96
53) 2,4-Dinitrophenol	14.957	184	14569	19.142 ng/ul	93
55) 4-Nitrophenol	15.057	109	22447	24.684 ng/ul	91
56) Dibenzofuran	15.227	168	178428	23.249 ng/ul	100
57) 2,4-Dinitrotoluene	15.198	165	48278	24.549 ng/ul	90
58) 2,3,4,6-Tetrachlorophenol	15.456	232	34865	25.569 ng/ul	96
59) Diethylphthalate	15.627	149	165527	24.057 ng/ul	99
61) Fluorene	15.879	166	145666	23.695 ng/ul	98
62) 4-Chlorophenyl-phenyle	15.856	204	76771	23.173 ng/ul	98
63) 4-Nitroaniline	15.909	138	30577	23.086 ng/ul	94
66) 4,6-Dinitro-2-methylph	15.962	198	26455	24.862 ng/ul	96
67) N-Nitrosodiphenylamine	16.073	169	131481	25.686 ng/ul	97
68) 4-Bromophenyl-phenylether	16.755	248	48124	25.112 ng/ul	93
69) Hexachlorobenzene	16.878	284	48968	25.060 ng/ul	96
70) Atrazine	17.019	200	53144	24.703 ng/ul	99
71) Pentachlorophenol	17.237	266	17388	20.082 ng/ul	97
72) Phenanthrene	17.624	178	256372	25.968 ng/ul	99
74) Anthracene	17.712	178	241770	24.658 ng/ul	99
75) 1,2,3,4-Tetrachloroben	13.635	216	62646	24.020 ng/uL	96
76) Pentachlorobenzene	15.145	250	58860	24.221 ng/uL	99
77) Carbazole	17.989	167	220127	25.577 ng/ul	99
78) Di-n-butylphthalate	18.512	149	287697	25.925 ng/ul	99
80) Fluoranthene	19.622	202	316390	27.219 ng/ul	99
82) Pyrene	19.986	202	309919	27.256 ng/ul	97
<ul><li>83) Butylbenzylphthalate</li><li>84) 3,3'-Dichlorobenzidine</li></ul>	20.850	149	121973	25.803 ng/ul	94
· · · · · · · · · · · · · · · · · · ·	21.767	252	45284	12.435 ng/ul	98
<ul><li>85) Benzo(a)anthracene</li><li>86) Bis(2-ethylhexyl)phtha</li></ul>	21.860	228	282649	26.643 ng/ul	100
87) Chrysene	21.720	149	182900	26.888 ng/ul	98
89) Di-n-octyl phthalate	21.931	228	267437	26.242 ng/ul	100
90) Benzo(b)fluoranthene	22.983 24.193	149 252	297137 280872	26.339 ng/ul 26.727 ng/ul	100 99
91) Benzo(k)fluoranthene	24.264	252		25.403 ng/ul	
93) Benzo(a)pyrene	25.116	252	250515 260911	26.024 ng/ul	99 98
94) Indeno(1,2,3-cd)pyrene	29.193	276	295675	26.824 ng/ul 26.354 ng/ul	96
95) Dibenzo(a,h)anthracene	29.252	278	246431	25.891 ng/ul	96
96) Benzo(g,h,i)perylene	30.415	276	197401	20.913 ng/ul	93
,					

<sup>(#) =</sup> qualifier out of range (m) = manual integration (+) = signals summed